A BILL FOR AN ACT

RELATING TO THE UNIFORM CONTROLLED SUBSTANCES ACT.

listed in this section are included in schedule V.

BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF HAWAII:

- 1 SECTION 1. Section 329-22, Hawaii Revised Statutes, is
 2 amended to read as follows:
 3 "\$329-22 Schedule V. (a) The controlled substances
- (b) Narcotic drugs containing nonnarcotic active medicinal ingredients. Any compound, mixture, or preparation containing limited quantities of any of the following narcotic drugs, which also contains one or more nonnarcotic active medicinal ingredients in sufficient proportion to confer upon the compound, mixture, or
- preparation, valuable medicinal qualities other than those
 possessed by the narcotic drug alone:
- 12 (1) Not more than 200 milligrams of codeine, or any of its salts, per 100 milliliters or per 100 grams;
- 14 (2) Not more than 100 milligrams of dihydrocodeine, or any of its salts, per 100 milliliters or per 100 grams;
- 16 (3) Not more than 100 milligrams of ethylmorphine, or any of its salts, per 100 milliliters or per 100 grams;

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1	(4)	Not more than 2.5 milligrams of diphenoxylate and not
2		less than 25 micrograms of atropine sulfate per dosage
3		unit;
4	(5)	Not more than 100 milligrams of opium per 100
5		milliliters or per 100 grams; and
6	(6)	Not more than 0.5 milligram of difenoxin and not less
7		than 25 micrograms of atropine sulfate per dosage unit.
8	(c)	Stimulants. Unless specifically exempted or excluded
9	or unless	listed in another schedule, any material, compound,
10	mixture,	or preparation that contains any quantity of the
11	following	substances having a stimulant effect on the central
12	nervous sy	ystem, including its salts, isomers, and salts of
13	isomers.	
14	(d)	Depressants. Unless specifically exempted or excluded
15	or unless	listed in another schedule, any material, compound,
16	mixture,	or preparation that contains any quantity of the
17	following	substances having a depressant effect on the central
18 .	nervous sy	ystem, including its salts, isomers, and salts of
19	isomers:	
20	(1)	Lacosamide [(R)-2-acetoamido-N-benzyl-3-methoxy-
21		<pre>propionamide], (Vimpat);</pre>

1	(2)	Pregabalin [(S)-3-(aminomethyl)-5-methylhexanoic
2		acid]; and
3	(3)	Brivaracetam ((2S)-2-[(4R)-2-oxo-4-propylpyrrolidin-1-
4		yl]butanamide) (Other names: BRV; UCB-34714; Briviact)
5		and its salts.
6	(e)	Approved cannabidiol drugs. A drug product in
7	finished	dosage formulation that has been approved by the United
8	States Fo	od and Drug Administration that contains cannabidiol
9	(2-[1R-3-	methyl-6R-(1-methylethenyl)-2-cyclohexen-1-yl]-5-
10	pentyl-1,	3-benzenediol) derived from cannabis and no more than
11	0.1 per c	ent (w/w) residual tetrahydrocannabinols."
12	SECT	ION 2. Section 329-38, Hawaii Revised Statutes, is
13	amended b	y amending subsection (i) to read as follows:
14	"(i)	Prescriptions for controlled substances shall be
15	issued on	ly as follows:
16	(1)	All prescriptions for controlled substances shall
17		originate from within the State and be dated as of,
18		and signed on, the day when the prescriptions were
19		issued and shall contain:
20		(A) The first and last name and address of the
21		patient; and

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(B) The drug name, strength, dosage form, quantity prescribed, and directions for use. Where a prescription is for gamma hydroxybutyric acid, methadone, or buprenorphine, the practitioner shall record as part of the directions for use, the medical need of the patient for the prescription.

Except for electronic prescriptions, controlled substance prescriptions shall be no larger than eight and one-half inches by eleven inches and no smaller than three inches by four inches. A practitioner may sign a prescription in the same manner as the practitioner would sign a check or legal document (e.g., J.H. Smith or John H. Smith) and shall use both words and figures (e.g., alphabetically and numerically as indications of quantity, such as five (5)), to indicate the amount of controlled substance to be dispensed. Where an electronic prescription is permitted, either words or figures (e.g., alphabetically or numerically as indications of quantity, such as five or 5), to indicate the amount

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of controlled substance to be dispensed shall be
acceptable. Where an oral order or electronic
prescription is not permitted, prescriptions shall be
written with ink or indelible pencil or typed, shall
be manually signed by the practitioner, and shall
include the name, address, telephone number, and
registration number of the practitioner. The
prescriptions may be prepared by a secretary or agent
for the signature of the practitioner, but the
prescribing practitioner shall be responsible in case
the prescription does not conform in all essential
respects to this chapter and any rules adopted
pursuant to this chapter. In receiving an oral
prescription from a practitioner, a pharmacist shall
promptly reduce the oral prescription to writing,
which shall include the following information: the
drug name, strength, dosage form, quantity prescribed
in figures only, and directions for use; the date the
oral prescription was received; the full name, Drug
Enforcement Administration registration number, and
oral code number of the practitioner; and the name and

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address of the person for whom the controlled substance was prescribed or the name of the owner of the animal for which the controlled substance was prescribed.

A corresponding liability shall rest upon a pharmacist who fills a prescription not prepared in the form prescribed by this section. A pharmacist may add a patient's missing address or change a patient's address on all controlled substance prescriptions after verifying the patient's identification and noting the identification number on the back of the prescription document on file. The pharmacist shall not make changes to the patient's name, the controlled substance being prescribed, the quantity of the prescription, the practitioner's Drug Enforcement Administration number, the practitioner's name, the practitioner's electronic signature, or the practitioner's signature;

(2) An intern, resident, or foreign-trained physician, or a physician on the staff of a Department of Veterans

Affairs facility or other facility serving veterans,

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•		exempled from registration under this chapter, sharr
2		include on all prescriptions issued by the physician:
3		(A) The registration number of the hospital or other
4		institution; and
5		(B) The special internal code number assigned to the
6		physician by the hospital or other institution in
7	•	lieu of the registration number of the
8		practitioner required by this section.
9		The hospital or other institution shall forward a copy
10		of this special internal code number list to the
11		department as often as necessary to update the
12		department with any additions or deletions. Failure
13		to comply with this paragraph shall result in the
14		suspension of that facility's privilege to fill
15		controlled substance prescriptions at pharmacies
16		outside of the hospital or other institution. Each
17		written prescription shall have the name of the
18		physician stamped, typed, or hand-printed on it, as
19		well as the signature of the physician;
20	(3)	An official exempted from registration shall include
21		on all prescriptions issued by the official:

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1		(A)	The official's branch of service or agency (e.g.,
2			"U.S. Army" or "Public Health Service"); and
3		(B)	The official's service identification number, in
4			lieu of the registration number of the
5			practitioner required by this section. The
6			service identification number for a Public Health
7			Service employee shall be the employee's social
8			security or other government issued
9			identification number.
10		Each	prescription shall have the name of the officer
11		stam	ped, typed, or handprinted on it, as well as the
12		sign	ature of the officer; and
13	(4)	A ph	ysician assistant registered to prescribe
14		cont	rolled substances under the authorization of a
15		supe	rvising physician shall include on all controlled
16		subs	tance prescriptions issued:
17		(A)	The Drug Enforcement Administration registration
18			number of the supervising physician; and
19		(B)	The Drug Enforcement Administration registration
20			number of the physician assistant.

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1	Each written controlled substance prescription issued
2	shall include the printed, stamped, typed, or hand-
3	printed name, address, and phone number of both the
4	supervising physician and physician assistant, and
5	shall be signed by the physician assistant. The
6	medical record of each written controlled substance
7	prescription issued by a physician assistant shall be
8	reviewed and initialed by the physician assistant's
9	supervising physician within seven working days."
.0	SECTION 3. New statutory material is underscored.
1	SECTION 4. This Act shall take effect upon its approval.

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Report Title:

Uniform Controlled Substances Act; Electronic Prescription; Schedule V

Description:

Amends the Uniform Controlled Substance Act by updating Schedule V and amending requirements for electronic prescriptions, for consistency with federal law. (CD1)

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